#### PATENT SPECIFICATION

NO DEVAMINGS

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COMPLETE SPECIFICATION

## Improvements in or relating to the Combination of Hydrogen and Oxygen

SPECIFICATION NO. 858,079

By a direction given under Section 17 (1) of the Patents Act 1949 this application proceeded in the name of Engelhard Industries, Inc., a corporation organised and existing under the Laws of the State of Delawsre, United States of America, of 113, Astor Street, Newark 2, New Jersey, United States of America.

THE PATENT OFFICE,

DS 87415/1(50)/R.153 200 2/61 PL

onthing hydrogen and only these containing water vapor or steam. These catalysts retain their initial high activity, and to steam. The term "platinum group metal" is used as meaning platinum, palladium, indium, rhodium, osmium or ruthenium, inidium, rhodium, osmium or ruthenium.

The catalyst surface area in excess of 30 slumins having a surface area in excess of 30 square metres per gram, and the silica content is, at least, 4 per cent by weight of the carrier. The catalytic metal may be present in the range of from 0.1 to 5 per cent by 70 weight of the combined catalytic metal and support, and of the various platinum group metals, e.g. platinum and incitals, palladium is preferred. If desired, and and rhodium, may be employed. The supportand in thodium, may be employed. The supportand the catalyst any be employed. The supportand the catalyst i.e. by treating the carrier or support manner, i.e. by treating the carrier or support with a solution of a suitable metal compound and then reducing the metal compound to said then reducing the metal compound to metal.

The temperature in the catalytic unit containing the catalyst may be in the range of

position of water in nuclear reactors.

bined to water, and reactors are often equipped tory, that the hydrogen and oxygen be recomreasons, it is desirable, and sometimes mandastruction at elevated temperatures. For these hydrogen and oxygen on materials of condifficulty resides in the corrosive effects of one which is potentially hazardous. A third times, converting an innocuous stream into oxygen concentration will be multiplied many the steam from a turbine, the hydrogen-For example, if steam is condensed, such as where these concentrations are low, the explosion hazard must always be considered. if the hydrogen and oxygen concentrations become appreciable and even in situations such venting is a hazardous procedure. Another difficulty is that explosions may occur where the vapor contains radioactive materials, increases unless the system is vented and, 20 oxygen in streams derived from nuclear reactors is troublesome for a number of reactors, one of which is that the gas pressure The presence of uncombined hydrogen and

[Price 3s. 6d.]

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#### COMPLETE SPECIFICATION

# Improvements in or relating to the Combination of Hydrogen amprovements

with recombining equipment, much of which is catalytic.

Processes for the catalytic combination of

hydrogen and oxygen in essentially dry gases 45 are well known, and the preferred catalyst is usually palladium or platinum on alumina these catalysts are used in contact with gas streams of high water vapor content, and at 50 elevated temperatures, they lose activity and the strength thereof deteriorates.

In accordance with the present invention, it has been found that nickel, or a platinum group metal, either singly or in combination, supported on a silica-alumina carrier, is a particularly advantageous catalyst for recombining water vapor or ateam. These catalysts retain their initial high activity, and containing water vapor or ateam. These catalysts retain their initial high activity, and 60 catalysts retain their initial high activity.

inidium, rhodium, osmium or ruthenium.

The catalyst support employed is silicallumina having a surface area in excess of 30 square metres per gram, and the silica content is, at least, 4 per cent by weight of the carrier. The catalytic metal may be present in the range of from 0.1 to 5 per cent by weight of the combined catalytic metal and metals, palladium is preferred. If desired, a mixture of platium group metals, e.g. platium group metals, e.g. platium and the desired, a mixture of platium group metals, e.g. platium and the catalyst may be employed. The support ted catalyst may be prepared in any suitable mener, i.e. by treating the carrier or support with a solution of a suitable metal compound with a solution of a suitable metal compound and then reducing the metal compound to

The temperature in the catalytic unit containing the catalyst may be in the range of

We, EMGRIHARD HANOVIA, IMC., formerly EMGRIHARD IMUSTRIES, IMC., a Corporation duly organised under the laws of the State of Mew Jersey, United States of America, of 100, Chestnut Street, Newark 5, New Jersey, United States of America, do hereby declare the invention, for which we pray that a patent the invention, for which we pray that a patent may be granted to us, and the method by which it is to be performed, to be particularly described in and by the following statement:—

The invention relates to an improved catalytic process for combining hydrogen and oxygen in the presence of steam or water vapor and, more particularly, the invention relates to the chemical combination of hydrogen and oxygen derived from the decomposition of water in nuclear reactors.

bined to water, and reactors are often equipped tory, that the hydrogen and oxygen be recomstruction at elevated temperatures. For these reasons, it is desirable, and sometimes mandahydrogen and oxygen on materials of condifficulty resides in the corrosive effects of 35 one which is potentially hazardous. A third times, converting an innocuous stream into oxygen concentration will be multiplied many the steam from a turbine, the hydrogen-For example, if steam is condensed, such as explosion hazard must always be considered. ٠٠ 30 where these concentrations are low, the if the hydrogen and oxygen concentrations become appreciable and even in situations Another difficulty is that explosions may occur such venting is a hazardous procedure. where the vapor contains radioactive materials, increases unless the system is vented and, reasons, one of which is that the gas pressure reactors is troublesome for a number of 20 oxygen in streams derived from nuclear The presence of uncombined bydrogen and

[Price 3s. 6d.]

o. A catalytic process for comming nytheogen and oxygen in the presence of steam, substantially as described in Example II of HASELTINE, LAKE & CO., London, W.C.Z.

London, W.C.Z.
Agents for the Applicants 6. A catalytic process for combining hydro- 15 ture in the range of from 100 to 800° C. the catalytic metal is maintained at a tempera-5. A process according to Claim 1 in which

Agents for the Applicants.

and rhodium. 3. A process according to Claim 1 in which the catalytic metal is a mixture of platinum the catalytic metal is palladium. 2. A process according to Claim 1 in which maintained above the dew point of the

the catalytic metal constitutes from 0.1 to 5 10 per cent of the combined weight of the 4. A process according to Claim I in which

catalytic metal and support.

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The oxygen in the effluent stream was calresidual gas was swept, by means of deoxy-genated hydrogen, into a sensitive analyzer (Hersch meter) for oxygen determination. condensed at room temperature, and the of 450° C. The effluent from the reactor was at atmospheric pressure and at a temperature passed through the high silica glass reactor the stoichiometric equivalent of hydrogen, volumes per million volumes of oxygen and bination boiler-electrolyzer such that 400 grams per hour of steam, containing 228 therein. Current was passed through the comof a sodium hydroxide solution contained electrodes for the electrolytic decomposition connected to the same boiler, which contained silica glass reactor, the latter reactor being and 50 ml. thereof were charged to a high was removed from the stainless steel reactor, exposure under these conditions, the catalyst was 400 grams per hour. After 37 hours of electrically heated to superheat the steam to a temperature of 450° C.; the vapor flow rate bed at a pressure of 600 psig. The reactor was 70 pressure boiler was passed through the catalyst less steel reactor, and steam from a highin Example I above, was placed in the stainexposure test.
A 100 ml. charge of the catalyst prepared pressure of 600 psig. used in the steam

unchanged from the attrition rate of fresh weight in six hours, which is also essentially loss by attrition was about I per cent by 100 spheres. In another empirical test of strength, within experimental error, as for unexposed bonuqa mpich was essentially the same, crushing strength of the spheres was Io culated to be less than 0.5 part per million, this being the limit of detection. The average

tially pure alumina, tested under identical of 0.5 per cent by weight palladium on essenresults were found using a catalyst consisting 105 In contrast to these results, the following catalyst.

for the fresh catalyst. exposed catalyst, versus 5 per cent by weight was 28 per cent by weight in 4 hours for the for the fresh catalyst, and the attrition loss 115 sbruog 8.2 susiev sbruog 3.1 ssw dignerie were present in the inlet gas. The crushing the effluent gas was greater than 20 parts per million were 110 present in the inlet gas, and 2 parts per million in the effluent when 9 parts per million were the effluent when 9 parts per million were a parts per million in the effluent when 9 parts per million were appearable. conditions with those given above. Oxygen in

area in excess of 30 square metres per gram on a silica-alumina carrier having a surface 125 nickel and a platinum group metal supported metal selected from the group consisting of prises contacting the mixture with a catalytic oxygen in the presence of steam which com-I. A process for combining hydrogen and 120 WHAT WE CLAIM IS:—

weight, the catalytic metal temperature being and a silica content of at least 4 per cent by

EXAMPLE II - aniroldD IISCE 0.054 per cent Water solubles

The catalyst was then dried in an oven at a temperature of 100° C, and assayed as

side of the spheres, after which they were

formation of a black Pd deposit on the outand Ma formate solution and heated, with the

Specifications of this carrier are 85 % Al<sub>2</sub>O<sub>3</sub>, 2.0% Na<sub>2</sub>O<sub>2</sub>, 0.15% Fe, 6.3% SiO<sub>2</sub>, and 6.2% loss on ignition.

spheres were used as the catalyst carrier. A catalyst was prepared as follows:—Commercial & diameter silica-alu

EXAMPLE 1

The invention will be further illustrated by

quantitatively, leaving the unreacted excess of

treatment will remove the minor constituent in stoichiometric proportions, the catalytic tures where the two gases are not present

to values below the explosive limit. In mixlimit the hydrogen and oxygen concentrations

from catalyst considerations, safe practice will

accordance with the present invention. Aside hydrogen and oxygen, may also be treated in

percentages of water vapor, in addition to

proportions. Gas compositions containing high taining hydrogen and oxygen in stoichiometric steam, either saturated or superheated, con-

in accordance with the present invention are

to 20,000 standard volumes per volume per

volume of catalyst per hour, preferably 5,000

to 50,000 standard volumes of vapor per the catalyst may be in the range of from 10

the reaction pressure as long as the dew point than 3200 psig. There are no limitations upon

of subatmospheric to super-critical, i.e., greater

temperature range is about 100 to 500° C.

maintained above the dew point. The preferred

proviso that the catalyst temperature must be

from 100 to 800° C., with the additional

The reaction pressure may be in the range

limitation is observed.

The space velocity of the gases passed over

The gas compositions which may be treated

reference to the following specific examples:-

the other constituent in the stream.

The spheres were treated with Na.PdCl.

0.40 per cent

diameter silica-alumina

but at atmospheric pressure rather than the temperature as the exposure, i.e. 450° C., glass. The evaluation was made at the same the catalyst was evaluated for activity in a separate reactor, fabricated from high silica reactor itself effected some oxygen removal, of nuclear reactor application. Since the tions closely simulating those used in one type above, to high-pressure steam, under condi-A stainless steel apparatus was used to expose the catalyst prepared in Example 1

Palladium

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